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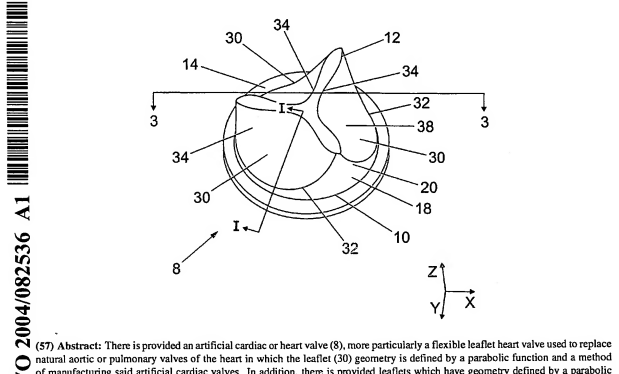
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[Continued on next page]

(54) Title: VALVE



natural aortic or pulmonary valves of the heart in which the leaflet (30) geometry is defined by a parabolic function and a method of manufacturing said artificial cardiac valves. In addition, there is provided leaflets which have geometry defined by a parabolic function.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

1	
2	"Valve"
3	
4	The present invention relates to artificial cardiac
5	or heart valves, more particularly to flexible
6	leaflet heart valves which are used to replace
7	natural aortic or pulmonary valves of the heart.
8	
9	Ideally artificial heart valves should work in a
10	similar fashion to natural heart valves in that when
11	blood flows in a particular direction the valve
12	adopts an open position to permit blood flow through
13	it, whereas when blood tries to flow in the opposite
14	direction the valve adopts a closed position
15	preventing the flow of blood in the reverse
16	direction through the valve (regurgitation).
17 [.]	
18	Natural heart valves use thin flexible tissue
19	leaflets as the closing members. In the closed
20	position the leaflets are arranged such that each
21	leaflet contacts its neighbour. This arrangement
22	serves to close the valve and prevent the back flow
	•

2

of blood through the valve. In the open position 1 the leaflets separate from each other and move 2 radially towards the inner walls of the blood vessel 3 in which the valve is located. This open 4 5 configuration of the valve permits the flow of blood through the valve. 6 7 8 A number of artificial cardiac valves have been 9 produced which comprise leaflets which open and close in a similar fashion to natural valve 10 leaflets. However, although the artificial valves 11 12 work in a similar manner to the natural valves, the geometries of the leaflets differ due to the 13 properties of the materials used in the construction 14 of the synthetic heart valves. 15 16 A number of factors have to be considered when 17 designing artificial heart valves of similar design 18 to natural heart valves. These include the pressure 19 gradient required to open and close the leaflets of 20 the valve, regurgitation, blood handling and 21 durability of the valve. 22 23 The leaflets of both natural and synthetic heart 24 valves must be capable of withstanding a high back 25 pressure across the valve when they are in the 26 closed position, yet be capable of opening with a 27 minimum of pressure across the valve in the forward 28 direction of blood flow. 29 30 This is necessary to ensure correct operation of the 31 valve even when blood flow is low. Further the 32

3

valve should open quickly and as wide as possible 1 when blood flows in the desired direction. The 2 maximum orifice of the valve in the open position is 3 generally dictated by the width of the valve. 4 5 In order to minimise closing regurgitation (reverse 6 blood flow through the closing valve) in the closed 7 position of the valve, the free edges of the 8 leaflets should come together to form a seal to 9 minimise the reverse flow of blood. 10 11 The valve design and the materials used for valve 12 construction should minimise the activation of both 13 the coagulation system and platelets. The flow of 14 blood through the valve should avoid exposing blood 15 to either regions of high shear or relative stasis. 16 17 Conventional heart valves typically comprise an 18 annular frame disposed perpendicular to the blood 19 flow. The annular frame generally has three posts 20 extending in the downstream direction defining three 21 generally U-Shaped openings or scallops between the 22 posts. The leaflets are attached to the frame 23 between the posts along the edges of the scallops 24 and are unattached at the free edges of the leaflets 25 adjacent to the downstream ends of the posts. 26 27 International Patent Application WO 98/32400 28 entitled "Heart Valve Prosthesis" discloses a 29 cardiac valve design, using closed leaflet geometry, 30 comprising essentially a trileaflet valve with 31 leaflets moulded in a geometry derived from a sphere 32

1	towards the free edge and a cone towards the base of
2	the leaflets. The spherical surface, defined by its
3	radius, is intended to provide a tight seal when the
4	leaflets are under back pressure, with ready opening
5	provided by the conical segment, defined by its
6	half-angle, at the base of the leaflets.
7	
8	International Patent Application WO 01/41679
9	discloses a heart valve wherein the leaflets have
10	been designed to facilitate wash out of the whole
11	leaflet orifice including the area close to the
L2	frame posts. This application teaches that stresses
13	are highest in the region of the commisures where
L 4	loads are transmitted to the stent, but they are
L5	reduced when the belly of the leaflet is as low as
16	practicable in the closed valve. To ensure a belly
17	in the leaflet, the above application indicates that
L 8	there must be sufficient material in the leaflet.
L9	
20	In addition, in order to be suitable for
21	implantation, synthetic valves should be
22	sufficiently durable such that they are clinically
23	functional for at least 20 years. Durability of the
24	synthetic leaflets depends on the materials from
25	which the leaflets are constructed and also the
26	stresses to which the leaflets are subjected during
27	use. However, although improvements have been made
28	to cardiac valves over recent years, problems still
29	exist with artificial valves. Although several
30	materials have suitable hydrodynamic properties,
31	many valves constructed using materials with
32	apparently suitable hydrodynamic properties

5

1	nevertheless fail during use, due to fatigue caused
2	by the repeated stresses of cycling from a closed to
3	an open position.
4	
5	The present inventor have surprisingly found that,
6	by using leaflets with parabolic configuration in
7	cross section, stresses of the leaflets can be
8	reduced and hence the lifespan of the valve may be
9	improved.
10	
11	It is an aim of the present invention to provide an
12	improved cardiac valve prosthesis.
13	
14	Thus, according to the present invention, there is
15	provided a cardiac valve prosthesis comprising:
16	
17	a frame and at least two flexible leaflets;
18	
19	wherein the frame comprises an annular portion
20	which, in use, is disposed substantially
21	perpendicular to the blood flow, the frame
22	having first and second ends, one of the ends
23	defining at least two scalloped edge portions
24	separated and defined by at least two posts,
25	each leaflet being attached to the frame along
26	a scalloped edge portion and being movable
27	between an open and a closed position,
28	
29	each of the at least two leaflets having a
30	blood inlet side, a blood outlet side and at
31	least one free edge, the at least two leaflets
32	being in a closed position when fluid pressure

1	is applied to the outlet side such that the at
2	least one free edge of a first leaflet is urged
3	towards the at least one free edge of a second
4	or further leaflet, and the at least two
5	leaflets being in an open position when fluid
6	pressure is applied to the blood inlet side of
7	the at least two leaflets such that the at
8	least one free edge of the first leaflet is
9	urged away from the at least one free edge of
10	the second or further leaflet;
11	
12	wherein, in a first plane perpendicular to the
13	blood flow axis, the length of each leaflet in
14	a circumferential direction (XY) between the
15	posts at any position along the longitudinal
16	axis (Z) of a post is defined by a parabolic
17	function.
18	
19	It is understood that reference to a parabolic
20	function includes reference to any
21	pseudotrigonmetric, pseudoelliptical, smooth
22	function or table of values that describe a geometry
23	which is substantially parabolic.
24	
25	The use of a pseudo function to describe a parabolic
26	function will be clear to one skilled in the art.
27	
28	Preferably the parabolic function defining the
29	length of a leaflet in the circumferential direction
30	(XY) between the posts at any position along the
31	longitudinal axis (Z) of a post is defined by
32	

1
$$Y_z = \left(\frac{4R}{L_z^2}\right) x \cdot (L_z - x)$$

2

3 Wherein $Y_z = Y$ offset at a particular co-ordinate X and Z

S R = parabolic maximum

6 L_z = straight line distance between a

7 first post and a second post of the frame

8 at a height Z

x = distance from origin of post towards

10 another post

11

wherein the length of the parabola can be

13 determined by

14

15 Length =
$$\int_{0}^{1} \sqrt{1 + \left(\frac{dy}{dx}\right)^2} dx$$

16

28

17 Preferably at least one correction factor can be applied to the measured lengths of for example L_z 18 or R to take into account changes in the dimensions 19 of the frame or material of the leaflet during the 20 21 cycling of the cardiac valve between an open and 22 closed position. For example, such changes, in the 23 dimensions may be, but are not limited to, inward 24 movement of the posts of the prosthesis on closure of the valve, stretch in leaflet material on closure 25 of the valve, or movement in the notional point of 26 coincidence of the leaflets. It will be clear to 27

the skilled man how to determine the correction

1 factor required in view of the frame and leaflet 2 material selected. 3 4 Preferably the correction factor is positive, 5 negative or zero. 6 7 The materials chosen to form the frame and the 8 leaflets of the prosthesis and the design of the 9 frame will influence to what extent the prosthesis, 10 including both the frame and the leaflets, yields to 11 the forces to which the prosthesis is subjected 12 during valve closure and opening. For example, 13 typically, inward movement of the posts of the 14 prosthesis occurs on closure of the valve due to the 15 force of the backward flow of blood on the leaflet. 16 This typically occurs to a greater extent at the 17 tips of the posts than where the posts meet the 18 frame. A correction factor is preferably included 19 in the determination of the XY lengths of the 20 leaflet at each height in Z to compensate for this 21 movement in the frame. 22 23 Preferably the cardiac valve prosthesis of the first 24 aspect of the invention comprises three leaflets. 25 In an embodiment of the valve comprising three 26 leaflets, one end of the frame of the cardiac valve 27 28 prosthesis defines at least three scalloped edge 29 portions separated by at least three posts, wherein each leaflet is attached to the frame along a 30 31 corresponding scalloped edge portion. 32

1 In such embodiments, preferably the three posts are rotationally symmetrically distributed around the 2 circumference of the frame. 3 4 5 Preferably the frame is a collapsible stent. This 6 may be advantageous as a collapsible stent may be 7 delivered to a patient by percutaneous delivery. 8 a preferred embodiment of the valve wherein the 9 frame is a collapsible stent, the collapsible stent 10 may be moved from a collapsed to an erect position 11 using an inflatable balloon when the stent is 12 suitably located in the patient. 13 The inventor has provided an improved cardiac valve 14 15 prosthesis by determining an advantageous leaflet 16 geometry. Indeed, a leaflet having such geometry 17 comprises an independent aspect of the present 18 invention. 19 According to a second aspect of the invention there 20 is provided a valve leaflet for use in the valve 21 22 according to the first aspect of the invention, 23 wherein the length of the leaflet in a 24 circumferential direction (XY) between the lateral 25 edges at any position along the lateral edge for 26 attachment to the post is defined by a parabolic 27 function. 28 29 Preferably the valve leaflet is a cardiac valve 30 leaflet for use in a cardiac valve prosthesis, more 31 preferably the cardiac valve prosthesis of the first 32 aspect of the invention.

1 2

3 As discussed above a parabolic function includes any

4 pseudotrigonmetric, pseudoelliptical, smooth

5 function or table of values that describe a geometry

6 which is substantially parabolic.

7

8 Preferably the parabolic function defining the

9 length of a leaflet in the circumferential direction

10 (XY) between the posts at any position along the

11 longitudinal axis (Z) of a post is defined by

12

13
$$Y_z = \left(\frac{4R}{L_z^2}\right) x \cdot (L_z - x)$$

14

15 Wherein $Y_z = Y$ offset at a particular co-ordinate X

16 and Z

17 R = parabolic maximum

18 L_z = straight line distance between a

19 first post and a second post of the frame

20 at a height Z

x = distance from origin of post towards

22 another post

23

24 wherein the length of the parabola can be

25 determined by

26

27 Length =
$$\int_{0}^{1} \sqrt{1 + \left(\frac{dy}{dx}\right)^{2}} dx$$

28

29

1 Preferably at least one correction factor can be 2 applied to the measured lengths of for example L, 3 or R to take into account changes in the dimensions 4 of the frame or material of the leaflet during the 5 cycling of the cardiac valve between an open and 6 closed position. 7 8 Preferably the correction factor is a positive, 9 negative or zero. 10 11 The leaflets are preferably formed from any 12 biostable and biocompatible material. 13 14 Preferably the leaflets are formed from Elasteon. 15 16 Preferably the leaflet has different thicknesses along a cross section defined by the intersection of 17 18 a plane perpendicular to the blood flow axis. 19 20 More preferably the thickness of the cross section 21 of the leaflet in the XY plane, defined by the 22 intersection of a plane perpendicular to the blood 23 flow axis, changes gradually and substantially 24 continuously from a thickest portion where the 25 leaflet is conjoined to the frame to a thinnest 26 portion at the midpoint of the XY plane of the 27 leaflet. 28 29 The leaflets of a valve as described above have a 30 top and bottom. In a preferred embodiment, wherein 31 the valve is a cardiac valve prosthesis of the first 32 aspect of the invention, the bottom of the leaflet

12

1 is attached to the scalloped portion and the top of 2 the leaflet defines the free edge. 3 4 Preferably the free edge of the leaflet is shaped to increase the length of the free edge of the leaflet 5 6 relative to the length of the leaflet in the XY 7 direction. 8 9 A valve leaflet of the second aspect of the 10 invention may be manufactured as part of the valve prosthesis or may alternatively be formed 11 independently and then attached to the valve once 12 13 formed. 14 Typically changing the diameter of the valve or 15 16 height of the posts of the frame affects the calculation of leaflet geometry i.e. the length of 17 18 the leaflets in the XY direction required to obtain 19 suitable closure of the valve. Conventionally, geometric scaling is employed to determine the 20 leaflet geometry for different diameters of valves, 21 22 but this technique lacks accuracy. 23 An advantage of the parabolic function described 24 herein to determine the XY length of the leaflet of 25 a cardiac valve is that the function can be used 26 irrespective of valve diameter or the height of the 27 posts of the frame to determine suitable leaflet 28 29 geometry and do not require the use of geometric 30 scaling. 31

1 Therefore functions disclosed by the present 2 Application which describe length in the circumferential direction (XY) of a leaflet e.g. the 3 4 leaflet geometry optimised for a 27mm inside diameter of stent can be used to describe the length 5 6 in the circumferential direction (XY) leaflet geometry for a stent of different diameter e.g. 17mm 7 8 inside diameter stent. 9 10 This makes the design and manufacture of valves of different diameters which comprise the leaflets of 11 12 the second aspect of the invention more convenient. 13 14 Preferably the free edge of the leaflet is shaped 15 such that in the longitudinal direction (Z) the free 16 edge of at least one leaflet is parabolic. 1.7 18 The parabola can be in either direction. However if the parabola extends away from the frame preferably 19 the maximum height of the parabola is $0\mu m$ to $500\mu m$ 20 21 more preferably 0µm to 100µm, even more preferably 22 0μm to 50μm higher than the notional straight line 23 between the ends of the parabola. 24 25 More preferably the free edge of at least one 26 leaflet is parabolic in the longitudinal direction 27 toward the scalloped edge portion of the frame such 28 that the maximum depth of the parabola is between 29 50μm to 1000μm, more preferably 50μm to 500μm, even more preferably 50 mm to 100 mm lower than the 30

1 notional straight line between the ends of the 2 parabola. 3 4 The inventor has surprisingly shown that by making 5 the free edge of valve leaflets parabolic, the 6 stress and strain characteristics of the leaflet at 7 the free edge are improved. 8 In particular embodiments the parabolic shape of the 9 10 free edge may be produced by trimming of the free 11 edge. 12 13 The valve of the first aspect of the invention can 14 be manufactured by any suitable method as known in 15 the art for example by adapting the method as 16 disclosed in WO 01/41679 or WO 02/100301. During 17 manufacture of a cardiac valve prosthesis it is 18 preferable if the leaflets are cast in a shape which 19 minimises the stresses in the leaflet during cycling 20 of the valve between the open and closed position. 21 Preferably, the leaflets are formed in a neutral 22 position, not fully open or closed. In addition, as 23 will be appreciated by those skilled in the art, in 24 a fully closed position the free edge of the 25 leaflets will be touching or almost touching each 26 other making manufacture of the leaflet difficult. 27 Once the length in XY of the leaflet, in respect of 28 the frame at a height Z has been determined the cast 29 shape of the leaflet can be defined to allow 30 manufacture of the leaflet on a forming element. 31

1	A preferred method of manufacture of the leaflets of
2	the first aspect of the invention has been developed
3	by the inventor. Indeed this preferred method
4	provides a further independent aspect to the
5	invention.
6	
7	According to a third aspect there is provided a
8	method of manufacturing a cardiac valve prosthesis
9	wherein the method comprises;
10	
11	 providing a forming element having at least
12	two leaflet-forming surfaces wherein the
13	forming surfaces are such that the length in
14	the circumferential direction (XY) of the
15	leaflet-forming surface is defined by a
16	parabolic function,
17	- engaging the forming element with a frame,
18	- applying a coating over the frame and the
19	engaged forming element, the coating binding to
20	the frame, the coating over the leaflet-forming
21	surfaces forming at least two flexible
22	leaflets, the at least two flexible leaflets
23	having a length in the circumferential
24	direction (XY) defined by a parabolic function
25	and a surface contour such that when the first
26	leaflet is in a neutral position an
27	intersection of the first leaflet with at least
28	one plane perpendicular to the blood flow axis
29	forms a wave,
30	 disengaging the frame from the forming
31	element.
32	

1 The coating is preferably a synthetic polymer 2 material, more preferably a synthetic resin or 3 plastics material. 4 5 As indicated above, when casting the leaflets, it is 6 desirable to keep the leaflets in a neutral position 7 and not touching each other. This is achievable by 8 casting the leaflets in a wave configuration. The leaflets are in a neutral position intermediate to 9 the open and closed position in the absence of fluid 10 11 pressure being applied to the leaflets. 12 13 The shape of the leaflet forming surfaces on which 14 the leaflets are cast is preferably defined by a 15 wave function. The wave function is thus applied to 16 the leaflet(s) to aid production of the leaflets 17 whose length in an XY direction has been determined. 18 The shape of the leaflet forming surfaces on which 19 20 the leaflets are cast may be defined by a first wave 21 having a first frequency. The first wave may be a 22 sinusoidal wave. 23 Alternatively, the shape of the leaflet forming 24 25 surfaces on which the leaflets are cast may be defined by at least two waves of differing 26 27 frequencies, which together form a composite wave. 28 A composite wave can be more complicated than a 29 30 single wave function. This provides a greater range of leaflet cast shapes, wherein the XY lengths of 31 the leaflet at each height 2 is defined by a 32

1	parabolic function or the like, in which the
2	leaflets may be manufactured.
3	
4	Preferably the wave defining the leaflet forming
5	surfaces and thus the cast shape of a leaflet is
6	asymmetric about the vertical mid plane parallel to
7	and intersecting the blood flow axis of the leaflets
8	when in use.
9	
10	Alternatively, the wave defining the leaflet forming
11	surfaces and thus the cast shape of a leaflet is
12	asymmetric about the vertical mid plane parallel to
13	and intersecting the blood flow axis of the
14	leaflets.
15	
16	In preferred embodiments the method further
17	comprises trimming the free edge of at least one fo
18	the leaflets formed. In particularly preferred
19	embodiments the method further comprises trimming
20	the free edge to a parabolic shape.
21	
22	It is preferred that the frame comprises three
23	posts. Preferably the number of leaflet forming
24	surfaces is equal to the number of posts.
25	
26	In the method of the invention the coating may be
27	applied to the frame in any suitable way known in
28	the art, for example using dip moulding,
29	conventional injection moulding, reaction injection
30	moulding or compression moulding.
31	

1

18

Dip moulding can be used to form surgical implants

2 of relatively complex shapes. Typically dip moulding is achieved by dipping a forming element 3 into synthetic polymer material, which may include 4 5 polymer resin or plastic material, removing the 6 forming element from the synthetic polymer material 7 and allowing the resultant coating of synthetic 8 polymer material on the forming element to dry or 9 cure. The moulded article is then removed from the 10 forming element. 11 12 A disadvantage of conventional dip moulding, as 13 described above, is that during the moulding of 14 intricate shapes, bubbles of air frequently become 15 trapped in cavities or recesses of the mould template. These bubbles of air remain trapped in 16 17 the moulded article when the article is cured and 18 give rise to holes or pits in the moulded article 19 rendering the moulded article unsuitable for use. Another problem encountered is that of providing an 20 21 even coating for articles of complex geometry. 22 example, precision coating is essential for 23 producing surgical implants of intricate shapes such as prosthetic heart valves. In particular, the 24 25 problems of bubbles and applying an even coating are 26 encountered when more viscous moulding materials are 27 used for moulding. 28 29 These problems with dip moulding can be minimised by 30 using inverted dip moulding. 31

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1	The co	ating may be applied over the frame by a
2	method	of inverted dip moulding comprising the
3	steps:	•
4		
5	-	submerging a forming element in a moulding
6		solution;
7	-	inverting said forming element whilst in the
8		moulding solution; and
9	-	isolating the forming element from the
10		moulding solution so that the coating thus
11		formed on the forming element can be dried
12		or cured.
13		
14	Invers	ion of the forming element whilst in the
15	mouldi	ng solution reduces the number of bubbles
16	formed	in the coating. Furthermore, such apparatus
17	enable.	s more efficient use of moulding solution and
18	lends .	itself advantageously to batch processing.
19		
20	In emb	odiments in which inverted dip moulding is
21	used,	the method may comprise the steps of:
22	(i)	attaching a forming element to a platform;
23	(ii)	sealing a housing to said platform to form a
24		closed chamber;
25	(iii)	filling said closed chamber with moulding
26		solution until the forming element is
27		submerged;
28	(iv)	inverting said closed chamber;
29	(v)	isolating the coated forming element from
30		the moulding solution.
31		

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20

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1	The coated forming element can be isolated from the
2	moulding solution by either breaking the seal and
3	removing the platform, for example by raising the
4	platform and thus the forming element out of the
5	solution, or by draining the moulding solution from
6	the closed chamber via outlet means.
7	
8	An apparatus for use in aspects of this invention in
9	which inverted dip moulding is used comprises:
10	
11	- at least one platform adapted to hold at
12	least one forming element;
13	- at least one housing having an open end
14	adapted to fit over said at least one
15	forming element;
16	 sealing means for reversibly sealing said
17	housing to said platform to form a closed
18	chamber suitable for holding a moulding
19	solution;
20	 means for inverting said closed chamber;
21	 closeable inlet means for introducing a
22	moulding solution into the closed chamber;
23	and
24	- closeable outlet means for releasing a
25	moulding solution from the housing.
26	
27	In particular embodiments of the manufacture of the
28	cardiac valve leaflet, in particular, coating of the
29	frame to form the leaflets, inverted dip moulding
30	and cutting or trimming of the leaflets, the forming
31	element is comprised of at least two portions

1 wherein portions are releasably attached to each 2 other. 3 4 Preferably releasable attachment of the at least two 5 portions of the forming element is provided by a 6 screw. 7 8 In a particular embodiment a first portion of the 9 forming element is a cardiac valve frame mounting 10 portion and a second portion is a base portion. The 11 base portion may be releasably attachable to the 12 inverted dip moulding apparatus. 13 14 The coating may be heated prior and / or during 15 moulding to aid movement of the material around the 16 forming element. This may be achieved by for 17 example heating at least a part of the moulding 18 apparatus is heated such that it heats the moulding 19 solution. 20 21 Preferably the synthetic polymer material is 22 biostable and biocompatible. 23 24 More preferably the synthetic polymer material is 25 Elasteon. 26 27 As described above, the inventor has found that 28 providing a parabolic shape to the free edge is 29 advantageous. 30 31 The parabolic shape may be formed during the coating 32 process or alternatively subsequent to manufacture

1 of the leaflets. It has been found that it may be 2 advantageous to cut the leaflets after formation. 3 For example, as discussed above, it may be 4 advantageous to trim the free edge of a leaflet, 5 e.g. to form a parabolic shape. 6 7 To date, conventional blades have been used to cut 8 moulded devices such as cardiac valves and leaflets 9 formed from synthetic polymer material. However, 10 these conventional blades become blunted over a 11 relatively short period of time, leading to the 12 production of moulded devices with a poor surface 13 finish on the cut edge. 14 15 To provide a high quality finish to a cut edge of the leaflet with minimal disruptions to the cutting 16 17 process to replace cutting blades it has been 18 determined that an ultrasonic cutting device may be 19 used. 20 The leaflets may be cut using an ultrasonic cutting 21 22 device comprising 23 (i) an ultrasonic transducer; 24 (ii) an elongate blade; and 25 (iii) attachment means to enable detachable 26 attachment of the blade to the transducer so 27 that, in operation, the transducer causes 28 the blade to vibrate in a direction along 29 the longitudinal axis of the blade. 30 31 It has been found that, for a given ultrasonic 32 frequency, by altering the dimensions of an elongate

1 blade, optimal operation of the cutting device can 2 be achieved. Reducing the amplitude of vibrations 3 perpendicular to the plane of the blade results in a cleaner cut. It has been found that by having a 4 5 blade of this particular construction precise 6 cutting of synthetic polymer material, for example, 7 resin and plastics materials can be achieved. 8 cutting device of the present invention is 9 particularly suitable for cutting acetyls, 10 polyurethane and polymeric materials. 11 12 Preferably the blade has a width to length ratio of 13 between 0.1 to 0.4. By width means the width of the 14 widest part of the blade and by length is meant the 15 length of the longest part of the blade. 16 Preferably the elongate blade has a length in the 17 18 range of 20 to 30 mm, a thickness in the range of 19 0.5 to 2 mm and a width in the range of 2 to 10 mm. 20 More preferably the width of the blade is between 5 21 and 8 mm. 22 23 Preferably the ultrasonic transducer or motor 24 produces vibrational energy at a frequency of 15 Hz. 25 26 The blade is provided with a terminal end, which is 27 the end furthest away from the transducer, which terminal end may have a single cutting edge and this 28 may be rounded in shape. Preferably the blade has a 29 plurality of cutting edges. Preferably the blade 30 31 has cutting edges along its longitudinal sides which 32 form a point at the terminal end of the blade, for

1	example in an arrowhead configuration. Preferably,
2	the longitudinal sides are arcuate in shape. In one
3	embodiment the blade is needle-shaped. Preferably
4	the blade is symmetrical in shape about its
5	longitudinal axis.
6	
7	The blade may be constructed from any suitable
8	material such as stainless steel, mild steel or
9	ceramic material. Preferably the blade is
10	constructed from a ceramic material. This is
11	advantageous as ceramic material is harder than
12	steel and remains cooler during operation of the
13	cutting device as there is less heat transfer to the
14	blade.
15	
16	Preferably the cutting apparatus further comprises
17	(i) a stylus for guiding the blade of the
18	cutting device on the surface of the article
19	to be cut which stylus comprises a rotatable
20	ball bearing mounted on an arm; and
21	(ii) attachment means for attaching the stylus to
22	the ultrasonic cutting device.
23	
24	The stylus is positioned so that, in operation, the
25	ball bearing is in contact with the surface of the
26	article to be cut. Preferably the rotatable ball
27	bearing is positioned above, but not in contact
28	with, the terminal end of the blade. Preferably the
29	outer most part of the rotatable ball bearing does
30	not extend to the outermost tip of the terminal end
31	of the blade so that, while the ball bearing is in
32	contact with the article to be cut, the cutting edge

1	of the terminal end of the blade penetrates the
2	article by a constant predetermined amount. This
3	results in a consistent and precise cut with each
4	part of the article experiencing the same exposure
5	to the cutting edge of the blade.
6	
7	The attachment means for attaching the stylus to the
8	ultrasonic cutting device may form part of means for
9	mounting the cutting device on a mounting table. The
10	means for mounting the cutting device on a mounting
11	table may further comprise means such as a 3-axis
12	drive unit as known in the art in which each arm of
13	the drive unit can move linearly in three directions
14	perpendicular to each other such that the ultrasonic
15	cutting device can be suitably positioned relative
16	to the article to be cut.
17	
18	Preferably the article to be cut is mountable on the
19	drive unit, for example the forming element on which
20	the cardiac valve leaflet to be cut is formed may be
21	mountable on the drive unit.
22	
23	A cardiac valve leaflet may be cut using an
24	ultrasonic vibrating blade comprising the steps of,
25	(i) positioning a blade relative to the heart
26	valve leaflet to be cut;
27	<pre>(ii) vibrating the blade;</pre>
28	(iii) moving the heart valve leaflet to be cut
29	relative to the vibrating blade or
30	alternatively moving the vibrating blade
31	relative to the heart valve leaflet to be

1	cut so that the blade cuts the heart valve
2	leaflet to the required shape.
3	
4	The heart valve leaflet may be mountable on the
5	mounting table while it is on the forming element on
6	which it was moulded.
7	
8	As described herein, an advantage of the valve of
9	the first aspect of the invention is that stresses
10	experienced by the leaflets during the cycling from
11	the closed to the open positions are minimised.
12	
13	By minimising the stresses present in the leaflets
14	of the valve during cycling from the closed to the
15	open position and back to the closed position the
16	lifetime of the synthetic leaflets is likely to be
17	increased.
18	
19	The present inventor has determined that fatigue
20	failures of previous synthetic valve are due to
21	bending stresses. In particular, the inventor has
22	determined that bending stresses affect synthetic
23	polymer valve material differently to non-synthetic
24	valve material.
25	
26	Indeed, the present inventor has determined that by
27	considering the stresses and strains of the leaflets
28	during cycling of the valve an optimal leaflet
29	geometry can be determined. This principle may be
30	applied to the design of other valves.
31	

27

1	Accordingly, in a further independent aspect of the		
2	invention there is provided a method of designing a		
3	cardiac valve prosthesis comprising the steps,		
4			
5	a) providing a model of a heart valve		
6	comprising a frame and at least two flexible		
7	leaflets,		
8			
9	b) generating loads experienced by at least one		
10	cardiac valve leaflet in use and applying these		
11	to the model,		
12			
13	c) determining the stress distribution of the		
14	leaflet,		
15			
16	d) changing the circumferential length of the		
17	leaflet in XY for any position in Z,		
18			
19	e) determining the new stress distribution of		
20	the leaflet,		
21			
22	f) repeating steps D and E to minimise local		
23	stress concentrations in the leaflet.		
24			
25	In preferred embodiments of this aspect of the		
26	invention, the cardiac valve prosthesis is a cardiac		
27	valve prosthesis of a first aspect of the invention.		
28			
29	In a particularly preferred embodiment the model		
30	comprises three flexible leaflets.		
31	•		

1 Preferably the method further includes the step of 2 adjusting the model to account for factors which influence the stress distribution of the leaflet 3 4 during the cycling of the cardiac valve between an 5 open and closed position. 6 7 More preferably, where the leaflets are formed from 8 synthetic polymer material, the method further 9 includes the step of adjusting the model to account 10 for factors depending on the synthetic polymer material of the leaflet which influence the stress 11 12 distribution of the leaflet during the cycling of 13 the cardiac valve between an open and closed 14 position. 15 16 Preferably the length of the leaflet in the 17 circumferential direction (XY) between the posts at 18 any position along the longitudinal axis (Z) of a 19 post is defined by a parabolic function and at least one correction factor. Preferably the correction 20 21 factor is used to compensate for at least one of, 22 but not limited to; inward movement of the posts of 23 the prosthesis on closure of the valve, stretch in 24 leaflet material on closure of the valve, or movement in the notional point of coincidence of the 25 26 leaflets. 27 28 Such correction factors are advantageous as they allow the determination of the XY length of the 29 30 leaflet to take into account factors which effect 31 the XY length of the leaflets required for closure 32 of the valve. For example, inward movement of the

29

1	posts of the prosthesis occurs on closure of the
2	valve, due to the force of the backward flow of
3	blood on the leaflet. This typically occurs to a
4	greater extent at the tips of the posts than where
5	the posts meet the frame. By providing a correction
6	factor in the determination of the XY lengths of the
7	leaflet at each height in Z to compensate for this
8	movement the leaflet length can be determined to
9	minimise bending stresses, in particular buckling of
10	the leaflet.
11	
12	The free edge of the leaflet of the cardiac valve is
13	particularly subject to stress and strain.
14	
15	Preferably the method further comprises the step of
16	providing different shapes and lengths of the free
17	edge of a leaflet.
18	
19	This is advantageous as it enables the effect of
20	trimming the leaflet to particular shapes, for
21	example parabolic, to be determined.
22	
23	Preferred aspects of the invention apply to each of
24	the other aspects mutatis mutandis.
25	
26	An embodiment of the present invention will now be
27	described, by way of example only with reference to
28	the accompanying drawings wherein;
29	
30	Figure la is a plan view of a trileaflet heart
31	valve in the closed position;
32	

30

1	Figures 1b, 1c and 1d show plan views of heart
2	valves with 3, 4 or 5 posts in which full
3	closure of the valve is achieved;
4	
5	Figures 1e, 1f and 1g show plan views of 3, 4
6	and 5 posted heart valves in which the length
7	XY of the free edge of the leaflets is defined
8	by a parabolic function;
9	
10	Figure 2a is a perspective view of an
11	embodiment of a trileaflet heart valve of the
12	present invention in a semi-closed position;
13	
14	Figure 2b is a perspective view of a prior art
15	trileaflet heart valve in a semi-closed
16	position;
17	
18	Figure 3 is a plan view of an embodiment of a
19	trileaflet heart valve of the present invention
20	in a semi-closed position;
21	
22	Figure 4a is a plan view of a prior art
23	trileaflet heart valve in a fully open
24	position;
25	
26	Figure 4b is a plan view of a prior art
27	trileaflet heart valve as shown in figure 4a in
28	a fully closed position;
29	
30	Figure 4c is a plan view of an embodiment of a
31	trileaflet heart valve according to the present
32	invention in a fully open position;

1	
2	Figure 4d is a plan view of an embodiment of a
3	trileaflet heart valve according to the present
4	invention as shown in figure 4c in a fully
5	closed position;
6	
7	Figure 5a is a cross section of the valve as
8	shown in figure 2a along line 3-3;
9	
10	Figure 5b is a cross section of the prior art
11	valve as shown in figure 2b along line 3-3;
12	
13	Figure 5c is a cross section of a valve with a
14	sigmoidal shaped leaflet in Z;
15	
16	Figure 6 is a plan view illustration of an
17	embodiment of a trileaflet heart valve of the
18	present invention;
19	
20	Figure 7a shows a partial cross section of a
21	post of an embodiment of a trileaflet heart
22	valve of the present invention in the open
23	position (II) and the closed position (I) of
24	the valve;
25	
26	Figure 7b shows a partial cross section of an
27	embodiment of a leaflet of the present
28	invention along the vertical midplane in the
29	open position (II) and closed position (I) of
30	the valve;
31	

32

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1	Figure 7c shows a partial cross section of a
2	post of a prior art valve in the open position
3	(II) and closed position (I) of the valve;
4	
5	Figure 7d shows a partial cross section of a
6	leaflet of a prior art valve along the vertical
7	midplane in the open (II) and closed (I)
8	position of the valve;
9	
10	Figure 8a shows the principal stress envelope
11	present in a prior art heart valve leaflet;
12	
13	Figure 8b shows the strain energy release
14	present in a prior art heart valve leaflet in
15	the X axis from a closed to open position;
16	
17	Figure 8c shows the strain energy release
18	present in a prior art heart valve leaflet in
19	the Y axis from a closed to open position;
20	
21	Figure 8d shows the resultant strain energy
22	release present in a prior art heart valve
23	during cycling from a closed to open position;
24	
25	Figure 9a shows the principal stress envelope
26	present in an embodiment of a heart valve
27	according to the present invention;
28	
29	Figure 9b shows the strain energy release
30	present in an embodiment of a heart valve
31	according to the present invention in the ${\tt X}$
32	axis from a closed to open position;

1	
2	Figure 9c shows the strain energy release
3	present in an embodiment of a heart valve
4	leaflet according to the present invention in
5	the Y axis from a closed to open position;
6	
7	Figure 9d shows the resultant strain energy
8	release present in an embodiment of a heart
9	valve leaflet according to the present
10	invention during cycling from a closed to open
11	position;
12	
13	Figure 10 is an illustration of an embodiment
14	of one leaflet according to the present
15	invention;
16	
17	Figure 11 is a diagrammatic representation of a
18	prior art leaflet moving from a semi-closed (a)
19	to successively more open position (b) and (c)
20	to a fully open position (d) illustrating the
21	formation of a bubble or buckle;
22	
23	Figure 12 illustrates a shape of the leaflet
24	being defined by a first wave further to
25	determination of the circumferential length of
26	the leaflet;
27	
28	Figure 13 is a graph of Cardiac Output (1/min)
29	against mean Pressure Gradient (mmHg);
30	

Т	
2	Figure 14a shows a sectional view of an
3	inverted dipping apparatus prior to moulding;
4	
5	Figure 14b shows a sectional view of an
6	inverted dip moulding apparatus post moulding;
7	
8	Figure 14c shows a cross sectional view of a
9	forming element suitable for use in the
10	moulding apparatus of the present invention;
11	·
12	Figure 15 is a perspective view of an
13	ultrasonic cutting device mounted on a mounting
14	table;
15	
16	Figure 16 is a view of the cutting apparatus of
17	an ultrasonic cutting device;
18	
19	Figure 17 is a perspective view of an
20	ultrasonic cutting apparatus according without
21	a stylus; and
22	
23	Figure 18 is a side view of ultrasonic cutting
24	apparatus without a stylus.
25	
26	As previously discussed, a number of designs have
27	been suggested for use in cardiac heart valves to
28	ensure that the heart valves have sufficient leaflet
29	material such that the valve is capable of opening
30	as wide as possible to the maximum orifice of the
31	valve, and that such opening requires as little

1 energy as possible and further that regurgitation of 2 blood through the valve is minimised. 3 4 In order to minimise the regurgitation of blood it 5 has been suggested that the free edge of the valve 6 is spherical in geometry to ensure that the free 7 leaflet edges are able to come together and seal 8 against one another. 9 10 US Patent 5,500,016 discloses a leaflet defined by 11 the equation: 12 $z^{2} + y^{2} = 2RL (x-q) - \alpha (x-q)^{2}$ 13 14 15 to describe the geometry of the leaflets. As Z, 16 defines the shape of the leaflet in the blood flow 17 axis and as Z is defined as z^2 then a leaflet 18 defined by the above would have a spherical geometry in the axis parallel to blood flow. International 19 20 Patent Application WO 98/32400 discloses that 21 spherical surfaces at the leaflet edges seal more 22 effectively than planar or conical surfaces. 23 International Application WO 01/41679 discloses that 24 stresses are highest in the region of the comissures 25 where loads are transmitted to the stent, but they 26 are reduced when the belly of the leaflet is as low 27 as practicable in the closed valve. 28 29 In addition, International Application WO 98/32400 30 also suggests that it is advantageous to provide a 31 spherical portion of leaflet adjacent to the base of the leaflet as it confers advantages in the stress 32

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1 distribution when the valve is closed and pressure 2 is greater downstream than upstream. 3 4 Thus, the prior art teaches that leaflets of heart valves should have considerable excess material in 5 the vertical axis Z, parallel to the blood flow to 6 7 enable a suitable seal to be achieved at the free 8 edge of the leaflet and to reduce the stress present 9 in the leaflet during open and closing. 10 11 As shown in figure 1b, 1c and 1d, the use of a frame comprising 3, 4 or 5 posts induces different angles 12 13 θ in the valve leaflets, to ensure a close fitting 14 tight seal of the leaflets, which minimises 15 regurgitation of blood through the valve. As the 16 number of posts increases, the smaller the angle $\boldsymbol{\theta}$ 17 and the more bent the leaflets are at a particular 18 point. In cycling between the open and closed 19 position, the valve will undergo considerable 20 flexing, particularly at angle θ . The smaller the 21 angle θ , the greater the stress experienced by the 22 valve at this point and the more the likely the 23 valve is to fail due to stress. 24 25 The material properties of tissue, which has low 26 stress at low and moderate strain means tissue . 27 valves are more able to cope with such flexing than synthetic materials. Synthetic materials typically 28 29 have different stress to strain relationships than 30 tissue and higher stress is typically experienced by 31 these materials at low and moderate strains. 32 means that flexing is more likely to cause damage to

1 leaflets constructed from synthetic material than 2 tissue material. 3 Previous valve designs have been largely based on 4 5 tissue valves and have not taken account of the 6 different material properties of synthetic material, 7 particularly synthetic polymer material. 8 9 In contrast to previous designs and teaching 10 concerning valve construction, which was driven by 11 the supposed need to obtain a close fitting seal of the leaflets, particularly at the free edge, the 12 13 leaflets of the valves of the present invention were 14 designed to minimise the stress experienced by the 15 leaflet during cycling between the open and closed 16 position. 17 To reduce the sharp curvature, which promotes stress 18 points at specific points along the free edge, the 19 20 length of the free edge (XY) of the leaflet was determined using a parabolic function. 21 parabolic length of the free edge can be determined 22 23 by using the distances between the posts of the 24 frame where the free edge is conjoined to the posts 25 and the parabolic maximum. 26 27 As shown in figures 1e, 1f and 1g the use of a 28 parabolic shape at the free edge results in a 29 gentler curvature of the leaflets and enables the 30 length of the material along the free edge to be 31 determined from a knowledge of the frame dimensions. 32 However, this design, contrary to previous teaching,

1 does not necessarily allow close fitting to be 2 achieved between the leaflets at all points along 3 the free edge. However, surprisingly, the seal 4 obtained between the leaflets using a parabolic or 5 like function was found to be sufficient to minimise 6 regurgitation of blood through the valve to the 7 required degree for the valve to be effective. 8 9 The determination of the length XY at the free edge 10 of the leaflet is important to ensure that closure 11 of the leaflets is achieved and to minimise the 12 excess material of the leaflets at the free edge 13 such that the free edges of the leaflets do not fold 14 over each other in the closed position. 15 16 In addition to allowing determination of the length 17 of XY at the free edge of the valve, the present Application also allows determination of the XY 18 19 lengths of the leaflets at all points in Z by using 20 a parabolic function to determine the shape of the 21 leaflets at all points in Z. 22 23 As shown in figures 5a, 5b and 5c, in the closed 24 position, the leaflet can be substantially linear 25 (figure 5a), have excess material such that a belly 26 forms (figure 5b) or have reduced XY lengths of the 27 leaflet towards the base such that the leaflet forms 28 a generally sigmoidal shape (figure 5c). 29 figures 5b and 5c the XY lengths of the leaflet and 30 thus the leaflet shape would be determined using a 31 non-continuous function.

32

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The inventor has determined the belly in the valve 1 2 as shown in figure 5b would create increased stress 3 in the belly region. Further, it has been deterimed that, as illustrated in figure 5c, a reduction of 4 5 material in XY towards the base of the posts 6 promotes an increase in the stress concentration at 7 the portion of the leaflets towards the free edge. 8 9 By determining the lengths XY of the leaflet as a parabolic function or the like at each point in Z, 10 11 such that the XY lengths in Z vary as a continuous 12 function, localised stress concentrations can be 13 minimised and a more uniform stress distribution 14 across the leaflet achieved. 15 16 As shown in figure 1a and figure 2a, a preferred 17 embodiment of the heart valve prosthesis 8 of the present invention comprises a stent or frame 10 18 19 which is substantially cylindrical. The frame has a 20 first end 12 and second end 14. The first end 12 21 comprises three scalloped edge portions 16a, 16b and 22 16c separated by three posts 18, each post having a 23 tip 20. The cardiac valve further comprises three 24 leaflets 30. Each leaflet 30 has a fixed edge 32 25 joined to a respective scalloped edge 16a, 16b or 16c of the frame 10 and a free edge 34 which extends 26 27 substantially between the tips 20 of the posts 18. 28 29 The leaflets 30 are configured to be movable from an 30 open to a closed position and from a closed to open 31 position. In an aortic position (when the 32 prosthesis is positioned at the site of the aortic

1	valve), the leaflets 30 have a blood inlet side 36
2	and a blood outlet side 38 and are in the closed
3	position when fluid pressure is applied to the
4	outlet side 38 i.e. by the blood of the aortic
5	artery and in the open position when fluid pressure
6	is applied to the inlet side 36 i.e. by the blood of
7	the ventricle. The leaflets are in a neutral
8	position intermediate to the open and closed
9	position in the absence of fluid pressure being
LO	applied to the leaflets.
L1	
L2	Where the valve is being used in a mitral position,
L3	between the left atrium and left ventricle of the .
L 4	heart, the orientation of the valve is opposite to
L 5	that described above such that blood flow from the
L6	left atrium moves the leaflets to an open position,
L7	the leaflets opening towards the left ventricle to
18	allow blood to flow into the left ventricle. Back
19	pressure from blood flow from the left ventricle
20	towards the left atrium causes the mitral valve to
21	close to minimise regurgitation.
22	
23	In figure 5b which is a sectional view along line 3-
24	3 illustrating the closed position of a leaflet of a
25	valve of the prior art, a 'belly' portion 40 exists
26	in the mid portion of the leaflet. This 'belly'
27	portion between the free edge and the central
28	portion of the leaflet causes leaflets of the prior
29	art to have a double curvature, a curve in XY and a
30	curve in Z. Further, the 'belly' shape 40 causes
31	leaflets of the prior art to be almost concave in

1	shape when viewed in cross section along the
2	vertical midplane of the leaflet.
3	
4	As shown in figure 5a, which is a sectional view of
5	the valve of the present invention along line 3-3 as
6	shown in figure 2a, no 'belly' is present in the
7	leaflets and in Z the leaflet in the closed position
8	is substantially linear.
9	
10	The conventional design including a 'belly' portion
11	was previously favoured as it was thought to
12	maximise sealing of the valve at the free edge and
13	minimise regurgitation.
14	•
15	However, the double curvature, which comprises
16	curvature in XY plane and in Z plane results in
17	excess leaflet material at both the open and closed
18	position which promotes the formation of a bubble or
19	buckle 50 in the leaflet material (as shown in
20	figure 11) during movement from a closed to open
21	position.
22	
23	This excess material is shown most clearly by
24	comparing figure 7d which shows a cross section of
25	the valve along the vertical midplane (line I-I of
26	figure 2b) of the leaflet 30 parallel to the blood
27	flow axis in a prior art leaflet with figure 7b
28	which shows a cross section along the vertical
29	midplane (line I-I of figure 2a) of a leaflet of the
30	present invention. This comparison clearly shows
31	that the leaflet 30 of the valve of the present
32	invention does not display a belly region 40.

T	indeed the cross section shown in ligure 15		
2	indicates that the leaflet shape of the present		
3	invention is substantially linear in the vertical		
4	direction in both the open and closed valve		
5	positions.		
6			
7	To determine the circumferential length of material		
8	in XY to remove the 'belly' 40 observed in prior art		
9	leaflets, the length in the circumferential		
10	direction (XY) of the leaflet for any position in \boldsymbol{z}		
11	must be determined, which still allows suitable		
12	opening and closure of the valve.		
13			
14	As shown in figure 6 the material of the leaflet		
15	must extend between the posts 18 such that in a		
16	closed position the free edge of the leaflets 34		
17	come together at point 42 to minimise regurgitation		
18	of blood through the valve.		
19			
20	This circumferential length (XY) can be		
21	mathematically defined using a parabolic function.		
22			
23	Function of a parabola		
24	$Y_z = \left(\frac{4R}{L_z^2}\right) x \cdot (L_z - x)$		
25			
26	Wherein $Y_z = Y$ offset at a particular co-ordinate X		
27	and Z		
28	R = parabolic maximum		
29	L_z = straight line distance between a		
30	first post and a second post of the frame		
31	at a height Z		

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1	X = distance from origin of post towards
2	another post
3	-
4	To calculate the circumferential length (XY) at a
5	height point of the posts for a leaflet defined in
6	the circumferential (XY) direction by a parabolic
7	function the following function can be used:
8	j wat can be abea.
9	length of parabolic curve = $\int_{0}^{t} \sqrt{1 + \left(\frac{dy}{dx}\right)^{2}} dx$
10	
11	This allows a circumferential length (XY) to be
12	determined at each height point in Z.
13	
14	Thus as shown in figure 10 the circumferential
15	length (XY) can be determined at Z1, Z2, Z3Zn.
16	
17	The length of the leaflet in the circumferential
18	direction (XY) is calculated and repeated in the
19	radial direction (Z) to provide the complete
20	geometry of the leaflet.
21	
22	As the dimensions of the scallop edge 32 of the
23	frame 10 as defined by the posts 18 of the frame can
24	be determined by measuring the frame, then the
25	straight line distance between a first post and a
26	second post of the frame at a height Z (L_z) for a
27	leaflet 30 can be determined by measuring the
28	distance between the two posts 18 at several height
29	points in Z (where Z is a particular height along
30	the posts). This post to post distance can then be

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used in the equation detailed above to generate a 1 2 parabola (P) at each height point. In the embodiment shown, due to the scallop shape 32 3 defined by the posts 18 the circumferential length 4 of the leaflet in XY will decrease moving from the 5 6 first end at the tip 20 of the posts toward the 7 second end of the frame 14 at the base of the posts. 8 The more height points which are chosen, the more 9 lengths (P) which can be calculated along Z. If a large number of height points are chosen the lengths 10 determined by the parabolic function moving from the 11 12 tip of the posts to the base will vary in a substantially linear fashion. 13 14 The leaflets 30 of a valve 8 which are of 15 circumferential length (XY) as determined using the 16 above parabolic function will meet at the free edge 17 34 of the leaflet 30, but will not meet 18 significantly at points lower than the free edge 34. 19 The meeting of the leaflets at the free edge allows 20 regurgitation to be minimised without including 21 excess material or a belly region 40 in the leaflets 22 23 30. 24 The circumferential length (XY) can be further 25 adjusted to take account of factors which occur 26 during cycling of the heart valve. These factors 27 28 include inward movement of the posts 18 of the frame 10 due to pressure on the leaflets 30 during closing 29 30 of the valve. The amount of inward movement of the 31 posts 18 of the frame 10 is influenced by the 32 rigidity of the frame 10 and the pressure exerted on

45

the valve. The tips 20 of the posts 18 of the frame 1 2 10 move to a greater extent than the base of the 3 posts and as the scallop geometry between the posts 4 18 of the frame 10 is accurately known this differential movement can be taken into account when 5 determining the optimal circumferential length (P) 6 7 of XY in the leaflet 30. 8 9 In addition to the posts 18 of the frame 10 moving 10 toward each other during closure, the posts 18 also 11 move towards the centre point 42 where the leaflets 12 meet or the point of coincidence. 13 circumferential length XY of the leaflet can be 14 adjusted to account for this movement. 15 The material of the leaflet 30 typically has some 16 degree of elasticity and will stretch in response to 17 18 blood flow pressure. This stretching can again be taken into account in determining the lengths of the 19 20 leaflet 30 to ensure that a belly region 40 of the 21 valve is minimised. 22 23 As shown in figure 8a, analysis of the stresses over time incurred by heart valves during the cycling 24 25 process has revealed that the principal area of 26 stress 60 in existing cardiac valves is found close to the midpoint of the free edge of the leaflets. 27 28 29 Using the data from figure 8a, strain energy release in X and Y, as shown in figures 8b and 8c 30 31 respectively can be determined. Figure 8b shows 32 that leaflets of the prior art have a vertical

predisposition to defect propagation 62 at the free 1 2 edge 34. Figure 8c indicates that leaflets have a 3 predisposition to defect in the lateral dimension, 4 at an area 64 in the leaflet 30 lower than the free 5 edge of the leaflet 34, the lower area being located 6 above the central portion of the leaflet. In tests during cycling of cardiac valves it has been found 7 8 that over time, the stress in this lower area 9 promotes failure of defects in the material to 10 occur. These defects can cause valve failure. 11 12 The present invention has shown that analysis of the 13 dynamics of existing valves during the cycling 14 process has determined that the stress in this lower area is caused by the leaflets requiring to change 15 16 the direction of their surface curvature during 17 cycling. 18 19 In particular, as shown in figure 11, on cycling 20 from a closed to an open position a region lower 21 than the free edge forms a bubble like formation or 22 buckle 50 on the surface of the leaflet which is 23 opposite in direction to the curvature of the 24 surface of the rest of the leaflet. 25 26 On moving from the closed to open position, the bubble like formation 50 is forced to become 27 inverted such that it projects in an opposite 28 29 direction causing a whip like action in the leaflet 30 This whip like action promotes high stresses in 31 the area lower than the free edge 34 of the leaflet, 32 as shown in figures 8a, 8b, 8c and 8d.

1 2 The inventor has surprisingly determined, as shown 3 in figure 9a, that the principal stress envelope in relation to the valve as described in the present 4 5 application, wherein the circumferential length XY 6 of the leaflet at any point in Z is defined as a parabolic function, is decreased across the whole of 7 8 the valve. In particular strain energy release in X and Y, as shown in figures 9b and 9c respectively, 9 in relation to the valve of the present invention 10 indicates that a leaflet wherein the circumferential 11 12 lengths XY are determined by a parabolic function 13 has minimised predisposition to defect propagation 14 in the lateral dimension at an area in the leaflet 15 lower than the free edge of the leaflet and above 16 the central portion. 17 18 A reduction in the predisposition to defect propagation in the lateral dimension at an area in 19 20 the leaflet between the free edge of the leaflet and 21 the central portion in the leaflet of the present 22 invention is observed because there is less excess 23 material and thus minimal belly in the leaflet of 24 the present design. 25 26 On moving from the closed to open position a bubble like formation 50 is no longer created and thus a 27 28 whip like action does not occur in the leaflet. As 29 discussed, it is this whip like action which has been determined to promote high stresses in the area 30 lower than the free edge of the leaflet. As 31 32 illustrated by comparing figures 8a and 9a, in

1 contrast to the valves of the prior art, uniform 2 principle stress distribution, is observed across 3 the surface of the leaflet of the valve described in the present Application. 4 5 6 Minimisation of the regions of stress in the 7 leaflet, during cycling of the leaflet, will 8 increase the durability of the leaflet. 9. 10 Use of a parabolic function to determine the 11 circumferential lengths XY of the leaflet at each 12 height point in Z causes the vertical distribution 13 of lengths of the leaflet to be substantially linear 14 at the fully open and closed position. 15 16 As described above, it will be appreciated by those in the art that other functions with the addition of 17 18 suitable modifying factors could be used to derive a 19 function which substantially describes a parabola 20 and which leads to the vertical distribution of 21 lengths of the leaflet to be substantially linear at 22 the fully open and closed position, but which is 23 based on for instance an elliptical function. 24 25 As discussed, additional parameters may be included 26 in the parabolic function used to determine the 27 circumferential lengths XY of the leaflet. 28 additional factors may account for movement in the 29 posts of the stent, elasticity of the leaflet 30 material during movement of the leaflets from a 31 closed to an open position or other factors which

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1 occur during cycling which influence the length of 2 the leaflet require to allow closure. 3 4 The function described above explicitly determines 5 lateral lengths of the parabolic curve at any height 6 point in Z which is along a post of the frame. 7 view of this the above function can be applied to 8 any diameter of valve or valves with different 9 heights of posts, without the need for geometric scaling. This means that different dimensions of 10 11 valves can be manufactured with the same leaflet 12 geometry without further undue experimentation. 13 14 The surface contour of the leaflets 30 of the 15 embodiment described are such that in a fully open position, the intersection of the leaflets of the 16 valve perpendicular to the blood flow axis, forms a 17 18 substantially cylindrical shape. 19 20 In addition to the above, it has also been 21 determined that stress at the free edge of the 22 leaflet, as shown in figure 8a, can be further 23 reduced by trimming the free edge 34 of the leaflet 24 in the longitudinal direction (Z) such that the free edge is substantially parabolic 70, with the maximum 25 26 depth of the parabola being furthest from the 27 notional untrimmed free edge 74. The maximum depth of the parabola is generally located at the midpoint 28 29 of the free edge 72 (figure 9a). Figure 9a shows the effect of introducing a parabolic curve in the 30 31 vertical direction of the free edge. Comparison of 32 figures 8b, 8c and 8d with 9b, 9c and 9d shows that

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1 the strain energy release at the free edge is 2 significantly reduced through the introduction of 3 the parabola in the longitudinal direction (2). 4 5 Ideally the notional free edge 74 is trimmed in a 6 parabolic curve, wherein the maximum depth 72 of the 7 parabola 70 in the longitudinal direction toward the 8 second end of the frame is between 50μm to 1000μm, 9 more preferably 50 mm to 500 mm, even more preferably 10 50μm to 100μm lower than the notional straight line 11 74 between the ends of the parabola. 12 13 A different shape of cut, trim or notch can be 14 introduced in the free edge to decrease the stress 15 at the free edge. However, particular shapes of 16 cuts, trims or notches may introduce defects into 17 the leaflet which would decrease the leaflets 18 durability to stress. A parabolic trim as described 19 is therefore advantageous in that focal points of 20 stress are not introduced to the free edge of the 21 leaflet. Cuts, trims and notches which do not 22 create bending stresses at localised points on the 23 free edge are preferable. 24 25 In one embodiment a parabolic cut may be made using an ultrasonic cutting device. As shown in figure 1, 26 27 in one embodiment the ultrasonic cutting device 28 comprises an ultrasonic transducer (100); a blade 29 (110); and attachment means (120) to enable 30 detachable attachment of the cutting blade to the 31 transducer. The blade has two arcuate cutting edges

- 1 which meet at a point to form the terminal end of
- 2 the blade. In this embodiment the stylus is not
- 3 present. The ultrasonic cutting device is mounted on
- 4 the mounting table (130) by means of a clamping
- 5 assembly (140). The clamping assembly includes an
- 6 upright member (150) that extends from a first end
- 7 perpendicularly from the mounting table, a support
- 8 member (160) that extends laterally from the upright
- 9 member and is held relative to the upright member by
- 10 a fixing block (170), and a clamp (180) which
- 11 secures the ultrasonic cutting device to the clamp
- 12 support member. The clamp support member is
- 13 slideably moveable up and down a portion of the
- 14 upright member by turning of an adjusting screw
- 15 (190). In addition, the clamp support member is
- 16 slideably moveable laterally in relation to the
- 17 upright member, this movement being effected by the
- 18 rotation of a second adjusting screw (200). The
- 19 clamp support member is located between the fixing
- 20 block and a securing plate (210). The securing
- 21 plate can be moved towards the upright member to
- 22 secure the clamp support member at a suitable
- 23 position.
- 24 As shown in figure 16 an arm (220) can extend from
- 25 the clamp (180) to the cutting blade. A ball
- 26 bearing (222) is rotatably mounted at one end of the
- 27 arm and is positioned just above, but not in contact
- with, the blade. In use the ball bearing is in
- 29 contact with the surface of the article to be cut
- 30 and its position controls the extent of blade
- 31 penetration into the article.

1 Figure 17 shows a perspective view of the cutting 2 apparatus in position for operation without the 3 stylus guide. The heart valve leaflet to be cut is 4 mounted on a 3-axis drive unit (230). This drive 5 unit may be driven by electric motors. Figure 18 is 6 a side view of the embodiment shown in figure 17. 7 8 In the embodiment of Figures 17 and 18, movement of 9 the drive means causes the heart valve leaflet to be 10 cut to be brought into contact with the blade. 11 accurate positioning of the heart valve leaflet to 12 be cut, the cutting process may be accurately 13 repeated. A set pattern can then be followed and may 14 be instructed by a computer which drives the drive 15 means. 16 Leaflets of the geometry described herein can be 17 18 produced using methods known in the art such as injection moulding, reaction injection moulding, 19 20 compression moulding or dip moulding. 21 22 In one embodiment the heart valve leaflets may be 23 made by inverted dip moulding. As shown in figure 24 14a an embodiment of inverted dipping apparatus may 25 comprise a platform (1000) holding a forming element 26 (1110). A housing (1130) is sealed to the platform 27 to form a closed chamber (1140). The housing 28. comprises side walls (1150) and a ceiling (1160) and 29 is provided with inlet means (1170) which can be 30 closed by valve (1180). 31

32

1 The platform is adapted to hold at least one forming 2 Preferably the platform is adapted to hold 3 one forming element. By hold means the forming element is secured to the platform so that it will 4 5 remain in place even upon inversion or rotation of 6 the platform. Preferably the forming element is 7 releasably held on the platform. 8 9 The forming element has a shape so that when coated with the moulding solution it will produce an 10 article of the desired size and shape. The forming 11 element may comprise a core holding a frame which 12 when coated with the moulding solution will produce 13 a leaflet of the desired size and shape. 14 15 16 In a preferred embodiment, the forming element 17 (1110) is of two-part form, as is shown in Figure 18 14C. The forming element comprises a frame mount 19 (1112) fixed to a base portion (1114). A frame 8, for a heart valve prosthesis, can be mounted on the 20 frame mount 1112. The frame mount is fixed to the 21 base by fixing means for example a screw (1116) or 22 any suitable fixing means such as a bayonet fitting 23 24 or push fit fitting. The frame mount is removable 25 from the base portion. 26 27 A frame mount and base portion, (two part forming element) may be used during leaflet construction, 28 29 the frame mount being suitably shaped to a frame to 30 be mounted on the frame mount and allow the 31 ' production of the leaflets by dip moulding.

frame mount can also be used to hold the frame and

1 leaflets during subsequent cutting of the valve 2 The frame mount is releasably attachable 3 to the base forming element portion such that the 4 frame mount portion can be removed from the base portion so that the base portion may be reused. 5 6 frame mount portion may be releasably attachable to 7 the base portion by a screw. Should the frame mount 8 be damaged during the cutting stage the frame mount 9 can be discarded while retaining the base portion 10 and thus only a part and not the entire forming 11 element need be replaced. In addition, different 12 types of forming element mounts capable of mounting 13 frames of different diameters or with different 14 valve leaflet shapes can be fixed to the same base 15 portion thus reducing the need for complete forming 16 elements. 17 18 The housing (1140) has an open end (1142) so that 19 when placed on the platform (1000) the forming 20 element can extend into the housing. 21 The housing is of a shape and size so that it fits 22 23 over the forming element (1110) and has the capacity 24 to hold enough moulding solution to coat the forming 25 element. The housing has a ceiling (1160) which is 26 the part of the housing opposite to the platform. 27 The housing may have any suitable shape, for example 28 it may be a cylinder having one closed and one open 29 end, with its closed end being the ceiling. 30 31 Typically the platform and the housing are 32 constructed from steel.

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1			
2	The apparatus is provided with means for inverting		
3	the closed chamber. The inverted and open chamber		
4	is shown in figure 14b. Invertion of the housing		
5	may be provided by means for rotating the platform		
6	about a horizontal axis. In one embodiment, the		
7	platform is rotatable about a horizontal axis		
8	through the horizontal plane of the platform. This		
9	may be achieved by having the platform pivotally		
10	supported on a frame. The frame may comprise		
11	lateral pins which extend laterally from the frame		
12	into the platform so that the platform can rotate		
13	around them. In an alternative embodiment, the		
14	housing is rotatable about a horizontal axis in the		
15	horizontal plane of the open end of the housing.		
16	This may be achieved by having the housing pivotally		
17	supported on a frame. The frame may comprise		
18	lateral pins which extend laterally from the frame		
19	into the housing so that the housing can rotate		
20	around them.		
21			
22	Preferably inversion of the closed chamber is		
23	effected by drive means including a hand crank and		
24	an electric motor.		
25			
26	The closed chamber has closeable inlet means for		
27	introducing the moulding solution to the closed		
28	chamber. The inlet means may be closeable by means		
29	of a valve. The inlet means are preferably an		
30	opening in the ceiling of the housing and are		
31	provided with a pipe in connection with a central		
32	reservoir of moulding solution. In one embodiment		

30

1 the platform is provided with the inlet means. The 2 inlet means may alternatively be provided in one of 3 the side walls of the housing so that it will be in 4 a position close to the platform in the closed 5 chamber. In this embodiment the moulding solution 6 may be pumped from a reservoir into the closed 7 chamber via the inlet means. This latter embodiment 8 is preferred when more viscous moulding materials 9 are being used. 10 11 Preferably the inlet means and/or the outlet means 12 are heated. The moulding solutions generally used in the moulding of surgical implants are generally 13 14 viscous in nature and this viscous nature can make 15 the movement of the moulding solutions through the 16 inlet and outlet means difficult to achieve. Heating means can be incorporated in the moulding 17 18 apparatus and used to heat both the housing and the 19 inlet and outlet means. The raised temperatures of 20 the moulding solutions make these solutions less 21 viscous allowing easier movement of the solutions 22 through inlet and outlet tubes. 23 24 The housing has closeable outlet means. Preferably 25 an opening/pipe in the ceiling of the housing forms 26 the outlet means. When the housing is inverted then the moulding solution can be drained through such an 27 28 opening/pipe under the force of gravity. The outlet 29 means may be closeable by means of a valve.

1 Preferably, as in the embodiment shown in Figures 14a and 14b, the outlet means is also the inlet 2 3 means. 4 5 In operation, a forming element is releasably secured to the platform and a housing is placed over 6 7 the forming element and sealed to the platform. closed chamber thus formed should be in a position 8 whereby the forming element is upright. Moulding 9 solution is introduced into the chamber through the 10 inlet means until it reaches a level above the 11 12 forming element, e.g. level (1152) indicated in 13 Figure 14a. At this stage the inlet means is closed 14 by means of valve (1180). After a suitable period 15 of time, the platform, and thus the closed chamber, 16 is inverted by rotating, in this case, the platform 17 around a horizontal axis. The inverted chamber is 18 then left for a suitable period of time before the 19 housing/platform seal is broken and the housing is 20 lowered. This exposes the now-coated forming element in an inverted position. This can be seen 21 22 in Figure 14b. The moulding solution can then be 23 drained from the housing using the inlet means 24 (1170) which doubles as outlet means in this 25 embodiment. Alternatively the moulding solution can 26 be drained from the housing before the housing/platform seal is broken. The coating on the 27 forming element can now be dried/cured/treated 28 29 appropriately. 30 31 As the closed chamber is a sealed system it is possible to exchange the air present in the interior 32

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of the closed chamber, when moulding solution is not 1 present, with another solution or gas. The type of 2 3 solution or gas with which the mould chamber can be 4 filled prior to introduction of moulding solution 5 can be chosen in line with manufacturing 6 requirements. In this way, contact between the 7 mould solution and moisture in the air can be 8 avoided. 9 10 In one embodiment the apparatus comprises a 11 plurality of platforms and a plurality of housings. In this embodiment, preferably all the inlet means 12 13 are in connection with a central reservoir of 14 moulding solution, with the inlet means and the 15 reservoir forming a manifold. Preferably the manifold is heated. In this embodiment, preferably 16 17 all the platforms are pivotally supported as a unit 18 on a frame or all the housings are pivotally 19 supported as a unit on a frame. Batch moulding carries the advantages of having greater consistency 20 21 of results and of being more cost effective. 22 23 As discussed the circumferential length XY of the 24 leaflet at any height point in Z along the post of 25 the frame is explicitly provided by a parabolic 26 function or a pseudo function used to describe a 27 parabolic function. As is clear from figures 1e, 1f 28 and 1g, the manufacture of valve leaflets in the 29 closed position, as described herein, by dip 30 moulding or injection techniques would be difficult 31 as the free edges of the leaflets contact each 32 other. Although a forming element could be provided

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1 in which the valve leaflets were produced in the 2 open position, it is more desirable to form the 3 leaflet in a neutral position between the two 4 extremes of fully open or fully closed. 5 6 One method of forming the leaflets is to determine 7 the length of the leaflet in the XY direction for each point in Z for a preferred shape of leaflet. 8 9 10 On determining the length of the leaflet at each point in Z to minimise the formation of a belly in 11 the leaflet and using appropriate correction factors 12 to determine a final XY length at that point in Z, a 13 wave function can be applied to the leaflet at that 14 15 point in Z. As shown in figure 12 the wave function will change the shape of the leaflet at that point 16 in Z from a parabolic curve to a desired cast shape, 17 but the length of the leaflet as determined by the 18 initial parabolic shape will be maintained and 19 following manufacture of the valve, closure of the 20 21 valve, will cause the leaflet to adopt a parabolic 22 shape again at each point in Z. 23 24 The wave shape of the leaflet is used to provide a 25 forming element with leaflet forming surfaces of the 26 shape as defined by the waves arranged in Z for 27 casting of leaflets. 28 29 The valve is thus produced such that in a cast 30 position the leaflet is in neutral position, 31 intermediate the open and closed position in the 32 absence of fluid pressure being applied to the

1 leaflets. Production of the valve in the neutral 2 position means that the leaflets are substantially 3 free of bending stresses in this position. 4 5 The shape of the forming element, on which the 6 leaflet is formed, can be defined by one wave 7 function, or several wave functions which together 8 form a composite wave. 9 10 Regardless of the wave function used for the casting of the leaflet, the length of the leaflet is defined 11 at each point in Z along the post of the scallop by 12 a parabolic function or pseudo parabolic function as 13 14 described above together with any correction 15 factors. 16 The shape of the inner surface of the leaflets will 17 18 substantially replicate the shape of the forming 19 element. The shape of the outer surface of the 20 leaflets will be similar to the shape of the inner 21 surface, but variations will result e.g. from the 22 properties of the polymer solution and techniques 23 used to create the leaflet. 24 25 The leaflets of suitable length as defined by the 26 parabolic function and any correction factors and of 27 shape as defined by either a single or composite 28 wave function are attached to a suitable frame. 29 construction of a suitable frame will be obvious to 30 those skilled in the art. The frame can be made of 31 a biocompatible polymer, metal or composite.

1 frame can be coated with polyurethane to allow 2 integration of the leaflets. 3 Further to describing a first leaflet using the ٠4 5 above function, the remaining two leaflets of this 6 three leaflet embodiment can be determined by 7 rotating the geometry about the Z axis through 120° 8 and then through 240°. 9 10 Having formed the leaflets of the valve as described 11 above these can then be trimmed to introduce a 12 parabolic curve into the the edge of the leaflet not 13 attached to the frame (free edge) which extends 14 horizontally between two posts. The maximum depth 15 of the parabola being located between 50 µm to 1000 µm 16 lower than the notional straight line between the ends of the parabola toward the portion of the 17 18 leaflet which attaches the leaflet to the scallop 19 portion of the frame. 20 21 As shown in figure 13, surprisingly, in addition to 22 reducing the lateral stress of the valve, 23 determination of the length of the leaflet at each 24 point in Z according to a parabolic function not 25 only minimises the formation of a belly in the leaflet, but also reduces the pressure gradient 26 27 required to open the valve from a closed position. 28 29 The opening of a cardiac valve to as wide an orifice 30 as possible under minimal pressure gradients is a 31 key parameter in the design of synthetic heart 32 valves.

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A valve of the present invention may be used in any required position within the heart to control blood flow in one direction, or to control flow within any type of cardiac assist device.

Modifications and improvements can be incorporated without departing from the scope of the invention.

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1	Cla	ims
2		
3	1.	A cardiac valve prosthesis comprising:
4		
5		a frame and at least two flexible leaflets;
6		
7		wherein the frame comprises an annular portion
8		which, in use, is disposed substantially
9		perpendicular to the blood flow, the frame
10		having first and second ends, one of the ends
11		defining at least two scalloped edge portions
12		separated and defined by at least two posts,
13		each leaflet being attached to the frame along
14		a scalloped edge portion and being movable
15		between an open and a closed position,
16		
17		each of the at least two leaflets having a
18		blood inlet side, a blood outlet side and at
19		least one free edge, the at least two leaflets
20		being in a closed position when fluid pressure
21		is applied to the outlet side such that the at
22		least one free edge of a first leaflet is urged
23		towards the at least one free edge of a second
24		or further leaflet, and the at least two
25		leaflets being in an open position when fluid
26		pressure is applied to the blood inlet side of
27		the at least two leaflets such that the at
28		least one free edge of the first leaflet is
29		urged away from the at least one free edge of
30		the second or further leaflet;
31		

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T	wne	rein in a first plane perpendicular to the
2	blo	od flow axis the length of each leaflet in a
3	cir	cumferential direction (XY) between the
4	pos	ts at any position along the longitudinal
5	axi	s (Z) of a post is defined by a parabolic
6	func	ction.
7		
8		
9	2. The	cardiac valve prosthesis as claimed in
10	clas	im 1 wherein the parabolic function defining
11	the	length of a leaflet in the circumferential
12	dire	ection (XY) between the posts at any
13	posi	ttion along the longitudinal axis (Z) of a
14	post	is defined by
15		
16	$Y_z =$	$\left(\frac{4R}{L_z^2}\right)$ x. (L _z -x)
17		
18	Wherein	$Y_z = Y$ offset at a particular co-ordinate X
19		and Z
20		R = parabolic maximum
21		L_z = straight line distance between a
22		first post and a second post of the frame
23		at a height Z
24		x = distance from origin of post towards
25		another post
26		
27		and the length of the parabola defined by
28		the above is determined by
29		

1 Length =
$$\int_{0}^{1} \sqrt{1 + \left(\frac{dy}{dx}\right)^{2}} dx$$

2

4 3. The cardiac valve prosthesis as claimed in any preceding claim comprising three leaflets.

6

7 4. The cardiac valve prosthesis as claimed in any 8 preceding claim wherein the frame is a 9 collapsible stent.

10

11 5. The cardiac valve prosthesis as claimed in any
12 preceding claim wherein at least one leaflet is
13 configured to increase the length of the free
14 edge of the leaflet relative to the length of
15 the leaflet in the XY direction.

16

17 6. The cardiac valve prosthesis as claimed in
18 claim 5 wherein the free edge of the leaflet is
19 configured such that in a longitudinal
20 direction (Z) perpendicular to the XY direction
21 the free edge of the leaflet is parabolic.

22

23 A valve leaflet for use in the valve according 7. 24 to any one of claims 1 to 6, wherein said 25 leaflet has first and second lateral edges each 26 for attachment to a corresponding post, wherein the length of the leaflet in a circumferential 27 28 direction (XY) between the lateral edges at any position along the lateral edge for attachment 29 30 to the post is defined by a parabolic function.

29

1 A valve leaflet as claimed in claim 7 wherein 2 the parabolic function is defined by 3 $Y_z = \left(\frac{4R}{L_z^2}\right) x \cdot (L_z - x)$ 4 5 6 $Y_z = Y$ offset at a particular co-ordinate XWherein 7 and Z 8 R = parabolic maximum 9 L_z = straight line distance between a 10 first lateral edge for attachment to a 11 corresponding post and a second lateral 12 edge for attachment to second 13 corresponding post at a height Z 14 x = distance from origin of first 15 corresponding post towards second 16 corresponding post 17 18 and the length of the parabola defined by 19 the above is determined by 20 Length = $\int_{0}^{1} \sqrt{1 + \left(\frac{dy}{dx}\right)^{2}} dx$ 21 22 23 24 9. A method of manufacturing a cardiac valve 25 prosthesis wherein the method comprises; 26 (a) providing a forming element having at least 27 28 two leaflet-forming surfaces wherein the

forming surfaces are such that the length in

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1		the circumferential direction (XY) of the
2		leaflet-forming surface is defined by a
3		parabolic function,
4		(b) engaging the forming element with a frame,
5 .		(c) applying a coating over the frame and the
6		engaged forming element, the coating binding to
7		the frame, the coating over the leaflet-forming
8		surfaces forming at least two flexible
9		leaflets, the at least two flexible leaflets
10		having a length in the circumferential
11		direction (XY) defined by a parabolic function
12		and a surface contour such that, in use, when
13		the first leaflet is in the neutral position an
14		intersection of the first leaflet with at least
15		one plane perpendicular to the blood flow axis
16		forms a wave,
17		(d) disengaging the frame from the forming
18		element.
19		
20	10.	The method as claimed in claim 9 wherein the
21		valve is a valve according any of claims 1 to
22		6.
23		
24	11.	The method as claimed in claims 9 or 10 wherein
25		the forming element has three leaflet-forming
26		surfaces.
27		
28	12.	The method as claimed in any one of claims 9 to
29		11 further comprising the step of shaping a
30		free edge of a leaflet.
31		

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1	13.	The method according to claim 12 wherein said
2		free edge of the leaflet is shaped to a
3		parabola in a longitudinal direction (Z)
4		perpendicular to the XY direction.
5		
6	14.	A method of designing a cardiac valve
7		prosthesis of any of claims 1 to 6 comprising
8		the steps,
9		
10		a) providing a model of a heart valve
11		comprising a frame and at least two flexible
12		leaflets,
13		
14		b) generating loads experienced by at least one
15		cardiac valve leaflet in use and applying these
16		to the model,
17		
18		c) determining the stress distribution of the
19		leaflet,
20		
21		d) changing the circumferential length of the
22		leaflet in XY for any position in Z,
23		
24		e) determining the new stress distribution of
25		the leaflet,
26		
27		f) repeating steps D and E to minimise local
28		stress concentrations in the leaflet.
29		
30	15.	A method as claimed in claim 14 which further
31		includes the step of adjusting the model to
32		account for factors which influence the stress

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1	•	distribution of the leaflet during the cycling
2		of the cardiac valve between an open and closed
3		position.
4		
5	16.	A cardiac valve prosthesis as substantially
6		herein before described with reference to one
7		or more figures 2a, 3, 4a, 4b, 4c, 4d, 5a, 6,
8		7a, and 7b of the accompanying drawings.
9		•
10	17.	A leaflet as substantially herein before
L 1		described with reference to one or more figures
12		9a, 9b, 9c, 9d, 10 of the accompanying
L3		drawings.
L 4		

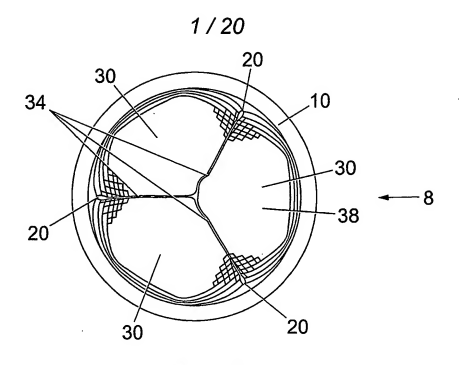
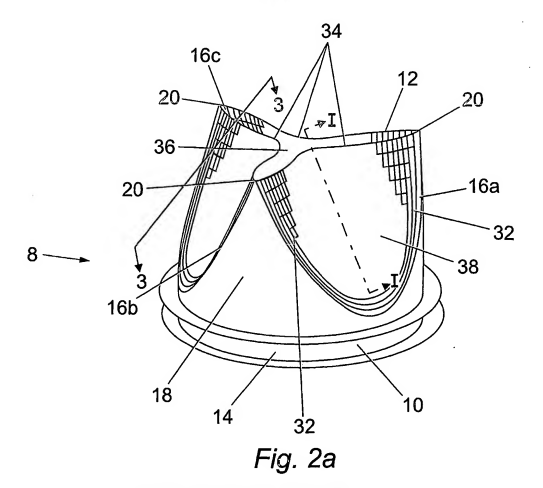


Fig. 1a



SUBSTITUTE SHEET (RULE 26)



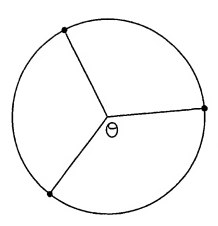


Fig. 1b

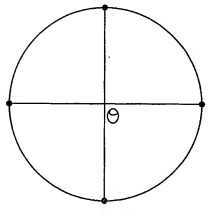


Fig. 1c

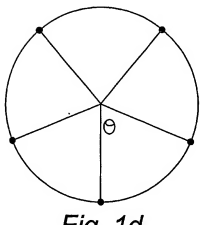


Fig. 1d

SUBSTITUTE SHEET (RULE 26)

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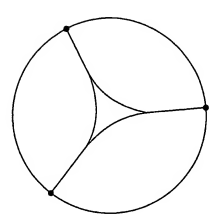


Fig. 1e

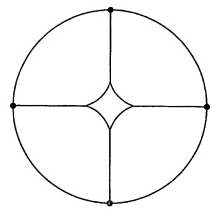


Fig. 1f

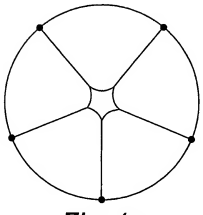


Fig. 1g

SUBSTITUTE SHEET (RULE 26)

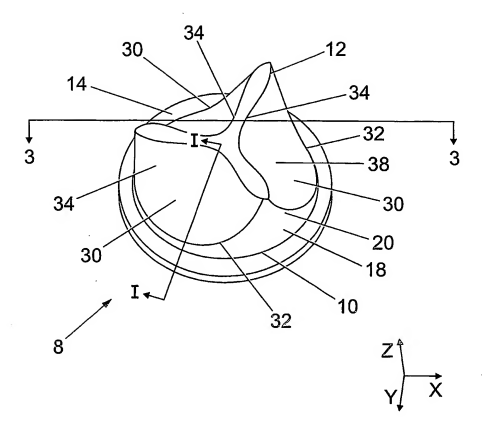


Fig. 2b

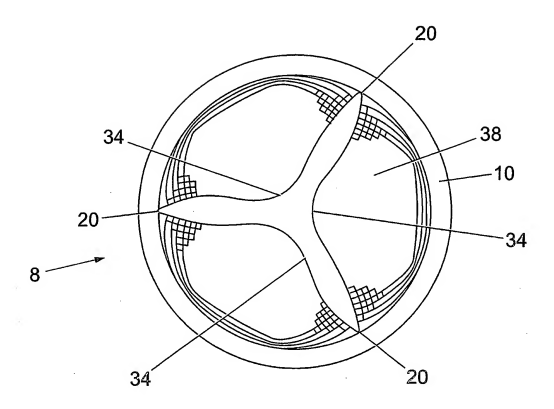
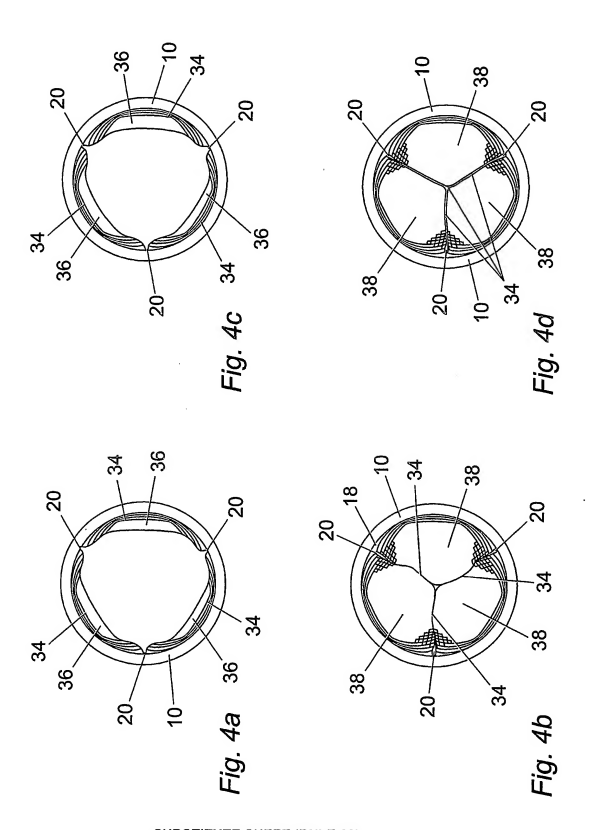
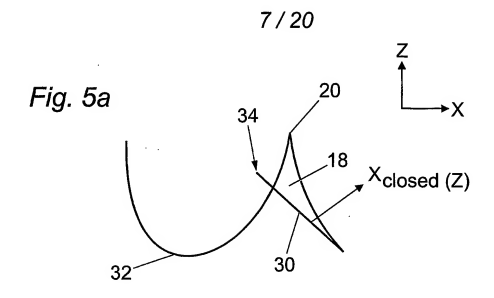
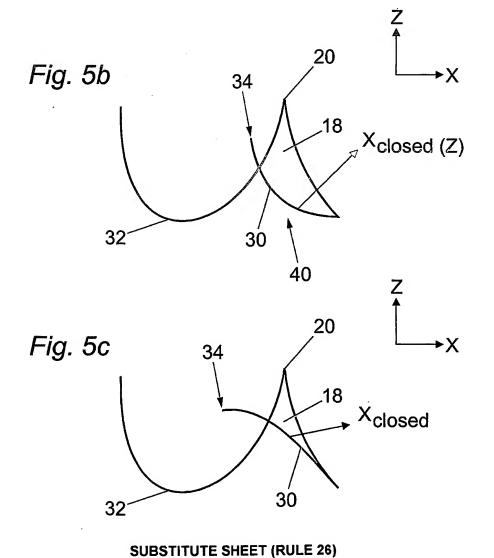


Fig. 3



SUBSTITUTE SHEET (RULE 26)





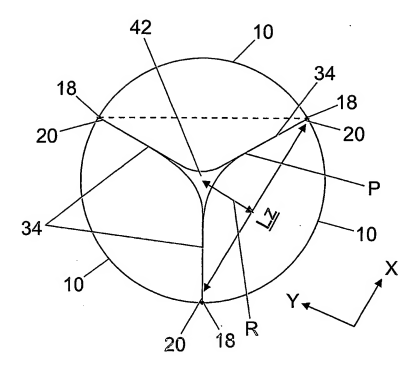
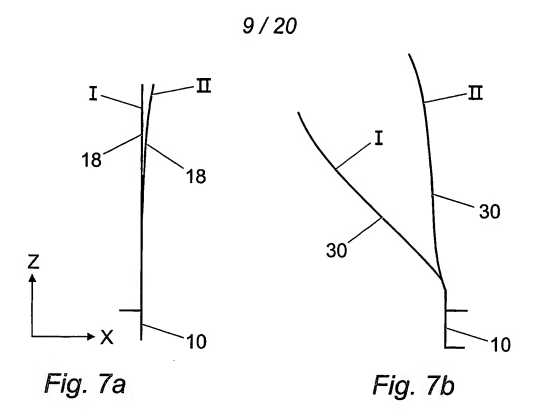
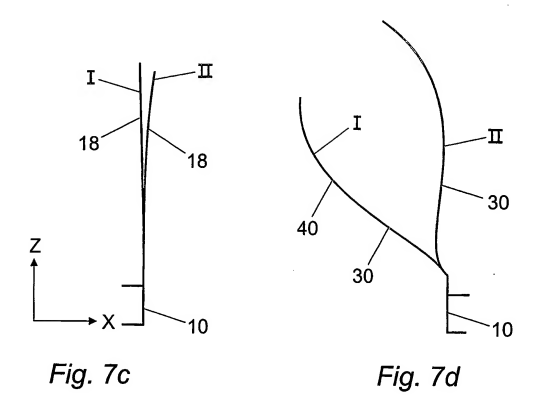
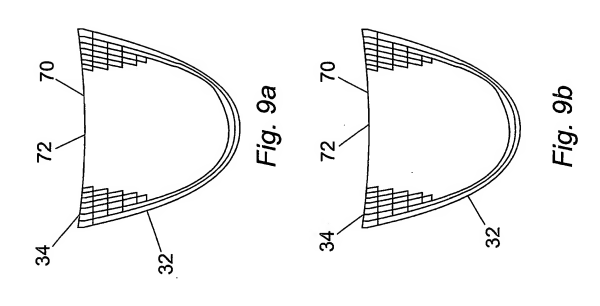


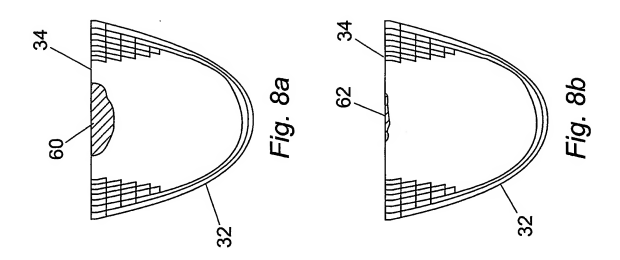
Fig. 6

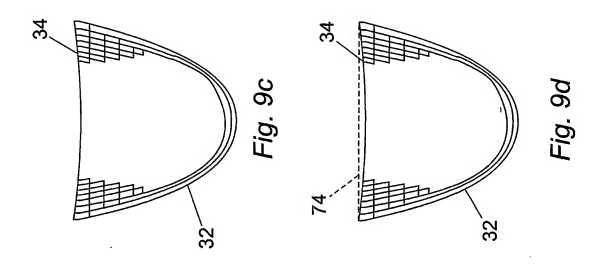


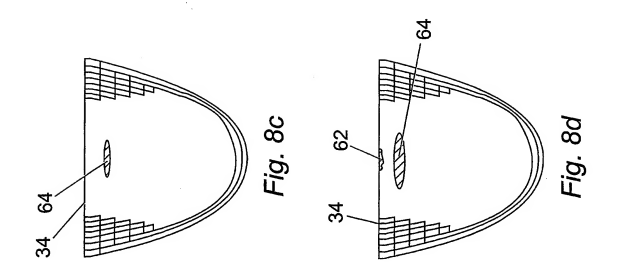


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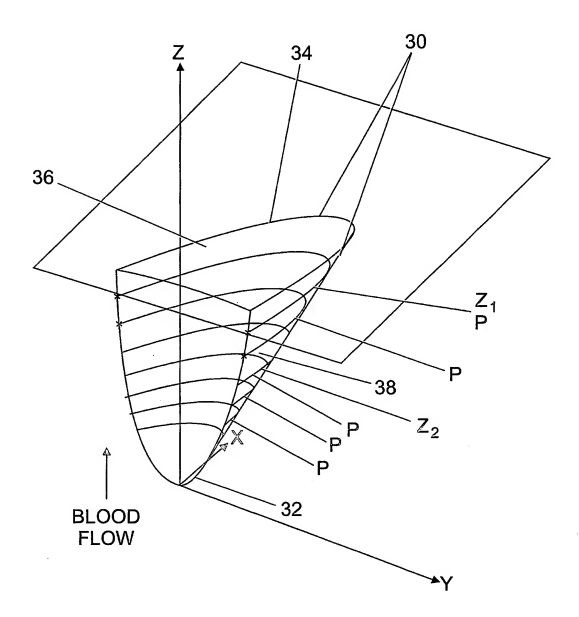
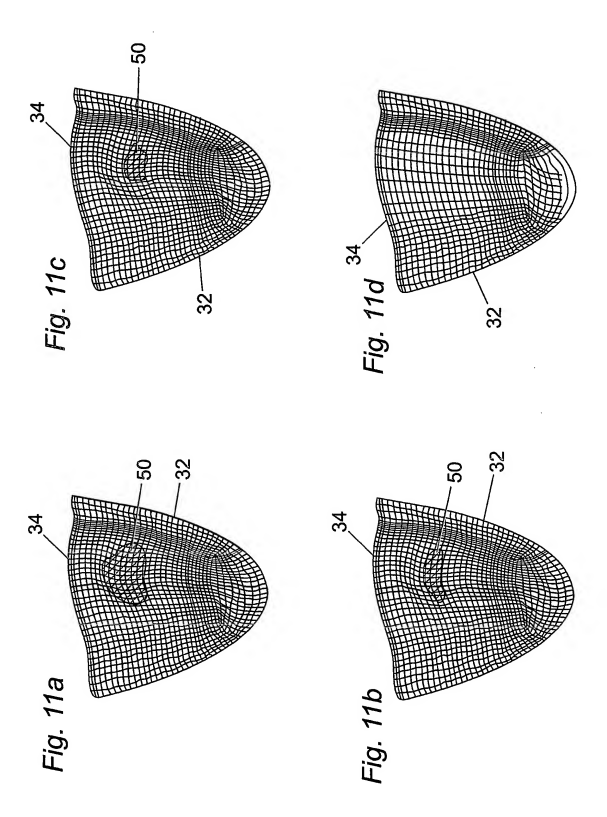


Fig. 10



SUBSTITUTE SHEET (RULE 26)

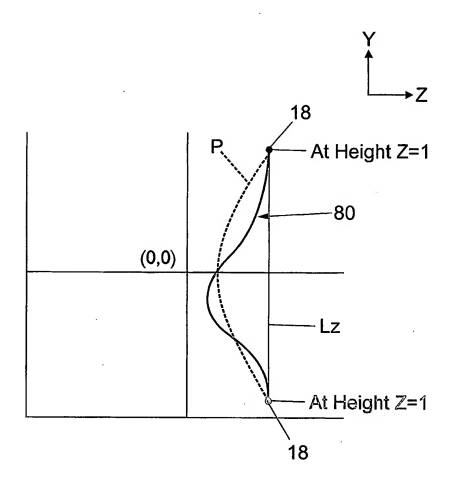


Fig. 12

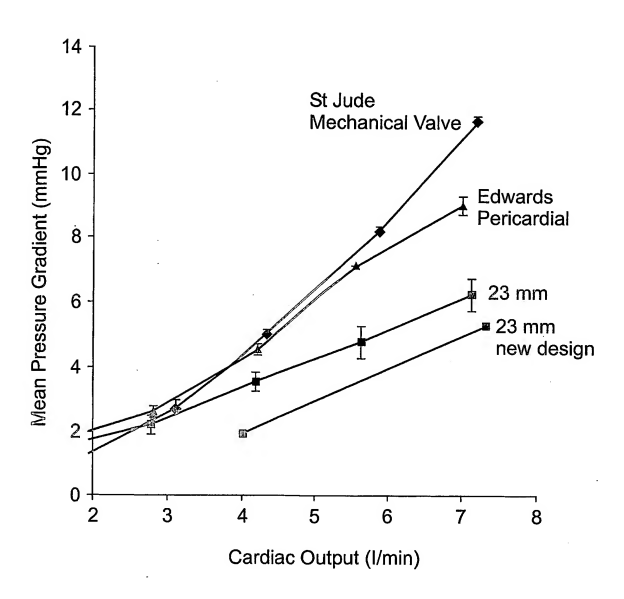
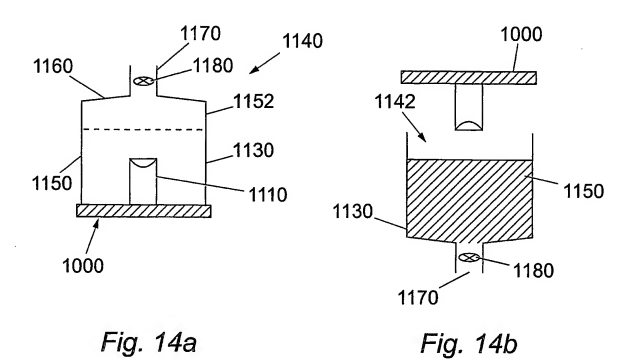


Fig. 13



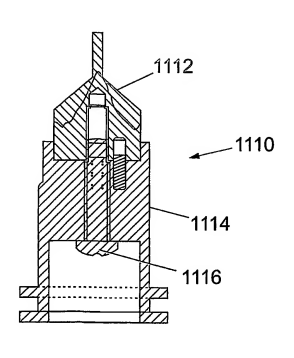


Fig. 14c

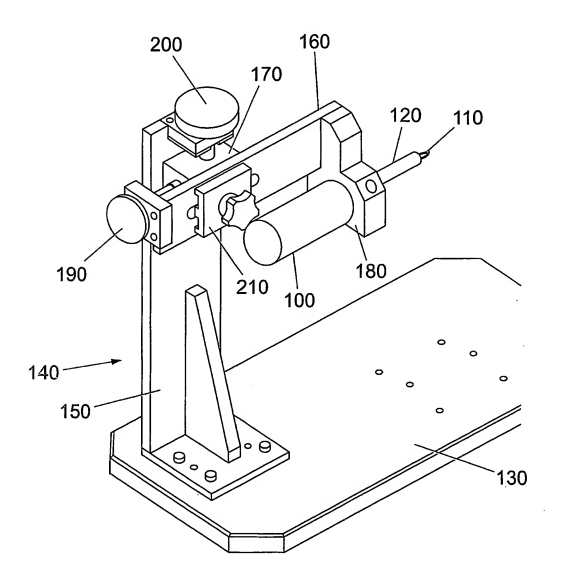


Fig. 15

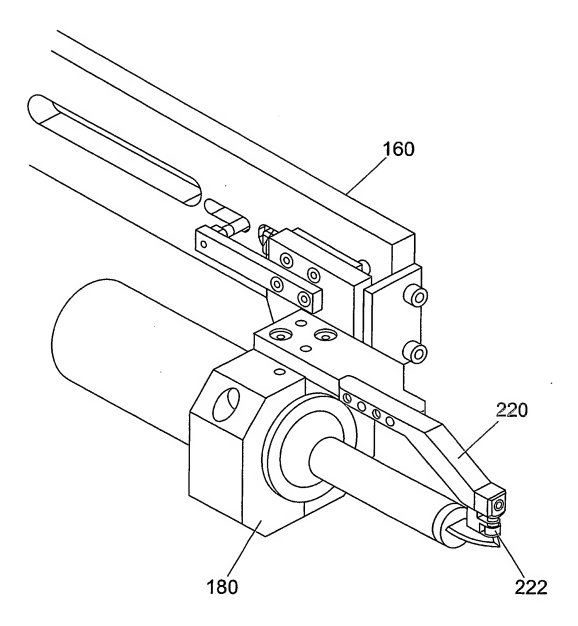


Fig. 16

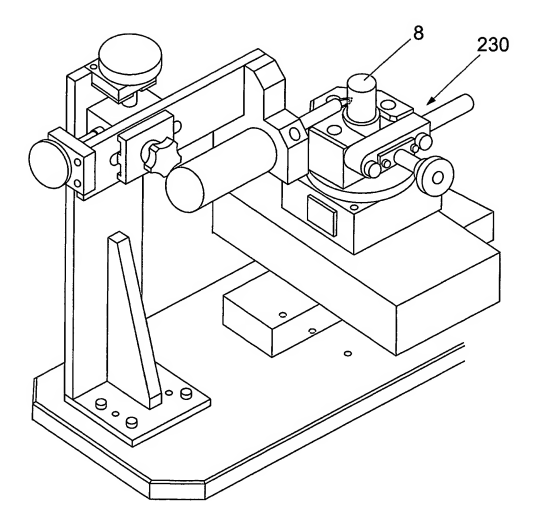


Fig. 17

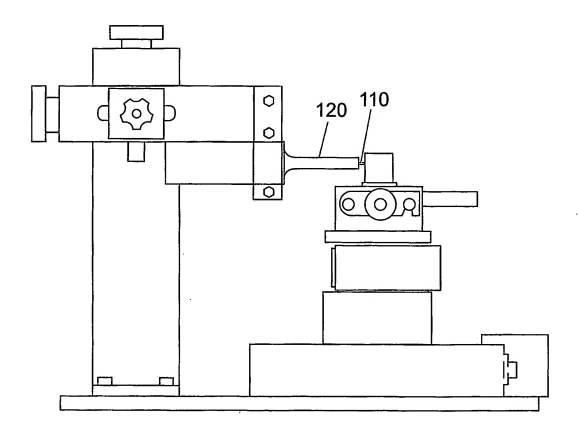


Fig. 18

INTERNATIONAL SEARCH REPORT

-----ational Application No

			.,				
A. CLASSI IPC 7	FICATION OF SUBJECT MATTER A61F2/24						
According to	International Patent Classification (IPC) or to both national classification	ation and IPC					
B. FIELDS	SEARCHED						
Minimum do	cumentation searched (classification system followed by classification $A61F$	on symbols)					
	lon searched other than minimum documentation to the extent that s	·					
Electronic d	ata base consulted during the international search (name of data bas	se and, where practical, sea	rch terms used)				
EPO-In	ternal						
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT						
Category *	Citation of document, with indication, where appropriate, of the rele	evant passages	Relevant to daim No.				
A	WO 01/41679 A (AORTECH EUROP LTD ; WHEATLEY 1-15 DAVID JOHN (GB); HAWORTH W S (GB); BER) 14 June 2001 (2001-06-14) cited in the application the whole document						
Α	WO 02/100301 A (AORTECH EUROP LTD) 19 December 2002 (2002-12-19) cited in the application the whole document						
A	WO 99/66863 A (SULZER CARBOMEDICS INC) 1-8 29 December 1999 (1999-12-29) the whole document						
P,X	US 6 613 086 B1 (MOE RIYAD E ET AL) 2 September 2003 (2003-09-02) column 5, line 54 - line 61; figure 18						
Funt	Further documents are listed in the continuation of box C. X Patent family members are listed in annex.						
° Special ca	pecial categories of cited documents:						
consid	"A" document defining the general state of the art which is not considered to be of particular relevance and the principle or theory underlying the invention						
filing d	invention invention invention invention invention connect be on particular relevance; the claimed invention document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is taken alone						
which i citation	is cited to establish the publication date of another nor other special reason (as specified)	"Y" document of particular re cannot be considered to	elevance; the claimed invention o involve an inventive step when the				
other n	O' document referring to an oral disclosure, use, exhibition or other means document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. P' document published prior to the international filling date but later than the priority date claimed "&" document member of the same patent family						
	actual completion of the international search		demational search report				
	30 June 2004 08/07/2004						
Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2							
	NL - 2280 HV Filjswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Newman, B					

INTERNATIONAL SEARCH REPORT

nternational application No. PCT/GB2004/001244

Box II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 16, 17 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: See FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)
This international Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Hemark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box II.2

Claims Nos.: 16, 17

There are no technical features in claims 16 and 17.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

INTERNATIONAL SEARCH REPORT

'Ional Application No 'GB2004/001244

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